Correlation and Prediction of the Transport Properties of Refrigerants Using Two Modified Rough Hard-Sphere Models¹

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ABSTRACT

Two methods are presented for the correlation and prediction of the viscosities and

thermal conductivities of refrigerants R11, R12, R22, R32, R124, R125, R134a, R141b

and R152 and their mixtures. The first (termed RHS1) is a modified rough hard sphere

method based on the smooth hard sphere correlations of Assael et al. The method

requires two or three parameters for characterizing each refrigerant, but is able to

correlate transport properties over wide ranges of pressure and temperature. The second

method (RHS2) is also a modified rough hard sphere method, but based on an effective

hard sphere diameter for Lennard-Jones (LJ) fluids. The LJ parameters and the effective

hard sphere diameter required in this method are determined from a knowledge of the

density - temperature behavior of the fluid at saturation. We show below that the RHS2

method can be used to correlate as well as predict the transport properties of refrigerants.

KEY WORDS: viscosity; thermal conductivity; rough hard sphere; refrigerants;

refrigerant mixtures;

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1. Introduction

Refrigerants and refrigerant mixtures are widely used as working fluids in many industrial applications, such as refrigerators, heat pumps and power plants. A knowledge of their transport properties is therefore of importance in the design and evaluation of these processes. Such knowledge is also of theoretical importance because it provides a framework for an understanding of intermolecular forces in refrigerant systems. Since it is unlikely that experimental measurements of transport properties at all conditions of interest can be found in the literature, reliable methods for their estimation are of considerable interest. One method that has been used successfully to correlate dense fluid transport properties was proposed by Assael et al. [1-5]. Their method (RHS3) is based on the rough hard sphere (RHS) theory, and employs a characteristic volume $V_{\mbox{\tiny o}}$ and coupling parameters (R_D , R_η , R_λ) for each substance to correlate self-diffusion, viscosity, and thermal conductivity over wide ranges of temperature and pressure. The method has been applied to n-alkanes [1], n-alkane mixtures [2], aromatic hydrocarbons [3], alkanols [4], and methane and ethane derived refrigerants [5]. A limitation of the method is that a complex series of calculations is required to obtain Vo and the coupling parameters. In this work, therefore, we propose a simplified and systematic way to evaluate these parameters and demonstrate the application of the technique (RHS1) to refrigerants.

A second method (RHS2) described below is based on our earlier work [6] on using the Lennard-Jones (LJ) fluid to obtain the hard sphere diameter σ_{HS} and hence the characteristic volume V_o required in the calculations. The LJ parameters σ_{LJ} and ϵ_{LJ} for each fluid are determined from a knowledge of the density-temperature behavior of the fluid at saturation.

We have used the two methods described above to correlate both high and low (saturated) pressure experimental data, and to compare the results with those of Assael et al. Special attention was given to the ability of the two methods to extrapolate data.

2. The Rough Hard Sphere Theory

The rough hard sphere concept was proposed by Chandler [7] and extended by Assael et al.[1-5] who showed that the reduced diffusivity D*, viscosity η * and thermal conductivity

 λ^* of all fluids can be expressed as universal functions of the reduced molar volume V_r (= V/V_o) as follows:

$$\log (D^*/R_D) = 3.285 - 31.74261 V_r^{-1} + 133.0472 V_r^{-2} - 285.1914 V_r^{-3} + 298.1413 V_r^{-4} - 125.2472 V_r^{-5}$$
(1)

 $\log (\eta^*/R_\eta) = 1.0945 - 9.26324 \ V_r^{-1} + 71.0385 \ V_r^{-2} - 301.9012 \ V_r^{-3} + 797.69 \ V_r^{-4} + 10.0385 \ V_r^{-2} - 301.9012 \ V_r^{-3} + 797.69 \ V_r^{-4} + 10.0385 \ V_r^{-2} - 301.9012 \ V_r^{-3} + 10.0385 \ V_r^{-3} + 10.0385 \ V_r^{-2} - 301.9012 \ V_r^{-3} + 10.0385 \ V_r^{-3} \ V_r^{$

$$-1221.977 V_r^{-5} + 987.5574 V_r^{-6} - 319.4636 V_r^{-7}$$
 (2)

$$\log \left(\frac{\lambda^*}{R_{\lambda}} \right) = 1.0655 - 3.538 \, V_r^{-1} + 12.120 \, V_r^{-2} - 12.469 \, V_r^{-3} + 4.562 \, V_r^{-4}$$
 (3)

where R_D , R_η and R_λ reflect the degree of coupling between translational and rotational motions of the molecules and account for deviation from the behavior of smooth hard spheres. In equations (1-3), the reduced diffusivity, viscosity and thermal conductivity are defined as

$$D^* = 5.030 \times 10^8 \,(M/RT)^{0.5} \,D \,V^{-1/3} \tag{4}$$

$$\eta^* = 6.035 \times 10^8 \, (1/MRT)^{0.5} \, \eta \, V^{2/3} \tag{5}$$

$$\lambda^* = 1.936 \times 10^7 \,(\text{M/RT})^{0.5} \,\lambda \,V^{2/3} \tag{6}$$

where M is the molecular weight, R is the gas constant, T is the temperature, η is the viscosity, λ is the thermal conductivity, and V is the molar volume of the substance (all properties being expressed in SI units).

The transport properties of a fluid at a given temperature and pressure can be calculated using equations (1-6), provided that the coupling parameters R_D , R_η and R_λ , and the characteristic volume V_o are known. Note that the volume V must also be available from experiment or from a pVT relation at these conditions.

3. Calculation of parameters

The parameters R_D , R_η , R_λ , and V_o are not equally significant at high pressures, as noted earlier by Dymond and Awan [8]. They found that the effect of nonspherical shape (and hence the coupling parameter) on diffusivity was negligible at high pressures, and that R_D could be set to unity. Similar behavior was found in this work in the case of viscosities of spherical fluids at high pressures, which could be correlated satisfactorily when R_η =1. On the other hand, diffusivities and viscosities were found to be very sensitive to changes in V_o . More emphasis was therefore placed on obtaining accurate values of V_o in the present

work. Also, calculated thermal conductivities showed only a slight dependence on V_o . This parameter was therefore estimated from viscosity data. (Note that diffusivities of refrigerants were not available, so that this property could not be used to obtain V_o).

The calculational procedure may be summarized as follows:

- 1. R_{η} was set equal to 1, or any other realistic value.
- 2. Experimental viscosities were used together with Eqs. 2 and 5 to calculate values of V_o , which are then fitted with a 4th order polynomial in temperature. (This requires an iterative procedure using the secant method).
- 3. The quantity $dy_1 = (\eta_{exp} \eta_{cal}) / \eta_{exp}$ was calculated
- 4. V_o and experimental data on thermal conductivity were used to calculate values of R_λ which were then fitted with a polynomial in temperature.
- 5. The term dy₂= $(\lambda_{exp} \lambda_{cal})/\lambda_{exp}$ was calculated.
- 6. Steps 1-5 were repeated until a minimum in $\Sigma dy_1^2 + \Sigma dy_2^2$ was obtained.

4. Application to Refrigerants

Four methane based refrigerants and five ethane based refrigerants were chosen for study because they had also been studied previously by Assael et al.[5]. Experimental viscosities of the nine refrigerants were obtained from the literature and consisted of 650 data at high pressures and 200 data at saturated pressure. The temperature range of the data was T/T_c =0.40 to 0.94 and the volume V at each temperature was obtained either from the literature or from the Tait equation reported by Assael et al. [5].

Table I lists the results of our calculations of the viscosities of the nine refrigerants. Average absolute deviations (AAD%) and maximum absolute deviations (MAD%) between experimental and calculated viscosities are listed for both the RHS1 method and the RHS3 method of Assael et al. In general, both methods show excellent agreement between calculated and experimental values. The large values of MAD for the RHS3 method are a result of the wider temperature range of the data used for comparison in the present work.

Experimental thermal conductivities of the nine refrigerants were also obtained from the literature and consisted of 550 data at high pressures and 75 data at low pressure.

The temperature range of the data was the same as for the viscosity. Table II lists the results of the comparisons for both the RHS1 and RHS3 methods. Again, the results show good agreement with each other and with experimental values. It is interesting to see that although V_o was obtained from viscosity data, calculated values of the thermal conductivity at both high and low pressures are still very satisfactory.

Finally, V_0 and R_{λ} were correlated as functions of temperature as follows:

$$V_o \times 10^6 = A_1 + A_2 T + A_3 T^2 \tag{7}$$

$$R_{\lambda} = B_1 + B_2 T + B_3 T^2 \tag{8}$$

where the coefficients A_i and B_i are listed in Table III.

5. Extension to Refrigerant Mixtures

The RHS1 method was extended to binary refrigerant mixtures using the mixing rules:

$$V_{o, m} = X_1 V_{o, 1} + X_2 V_{o, 2}$$
(9)

$$R_{\eta, m} = x_1 R_{\eta, 1} + x_2 R_{\eta, 2}$$
 (10)

$$R_{\lambda, m} = x_1 R_{\lambda, 1} + x_2 R_{\lambda, 2}$$
 (11)

where x_1 and x_2 are the mole fractions of component 1 and 2 and the subscript m denotes a mixture quantity. This mixing rule has no adjustable parameters, and does not require experimental data for the pure components. It has been used successfully by Assael et al. to calculate viscosities and thermal conductivities of n-alkane mixtures over wide ranges of temperature and pressure.

The results of the application of the mixing rule to refrigerant mixtures are given in Table IV. Both the RHS1 method and the RHS3 method were compared and found to work well for thermal conductivities, although large errors were observed for the RHS3 method in the case of R12+R22 because the temperatures of the data (200K) exceeded the fitting range of that correlation. Mixture viscosities, however, were not satisfactory. Both methods overpredicted the viscosities of R32+R134a mixtures by as much as 7% and underpredicted viscosities of R32+R124 by as much as 13%. This may be due to the fact that each component has a strong dipole moment (1.98 Debye for R32, 2.06 for R134a, and 1.47 for R124) so that a more realistic mixing rule may be required.

6. Calculation of V_o from the Lennard-Jones hard sphere diameter

Since the Lennard-Jones potential behaves like the hard-sphere (HS) potential for dense fluids, the hard-sphere diameter σ_{HS} can be obtained from the Lennard-Jones parameters σ_{LJ} and ϵ_{LJ} provided that a proper correlation between these quantities is established. The hard-sphere diameter σ_{HS} can then be used to determine the characteristic volume V_o as follows:

$$V_o = (\sigma_{HS}^3 / 2^{0.5}) N$$
 (12)

where the term in the brackets is the HS close-packed volume [9] of each molecule.

Given V_o , the coupling parameters R_D , R_η and R_λ can be obtained by fitting experimental data for the diffusivity, viscosity and thermal conductivity as described previously. We call the method with parameters obtained via the LJ hard sphere diameter the RHS2 method.

Several correlations for σ_{HS}/σ_{LJ} were examined. The most reliable was that proposed by Heyes [10] and given by:

$$\sigma_{HS}/\sigma_{LJ} = 1.0217 (1-0.0178 \text{ T}^{*-1.256}) / \text{T}^{* \text{ } 1/12}$$
 (14)

where $T^* = T/(\epsilon_{LJ}/k)$. This equation was obtained by fitting the simulation results for transport properties of LJ fluids, with the power of 1/12 coming from the scaling behavior of soft-sphere fluids.

LJ parameters for the nine refrigerants studied were determined from a LJ equation of state [11] at T/T_c =0.70 - 0.80 using saturated liquid density data. Two sets of data were used: densities from the Tait equation reported by Assael et al. and densities from NIST tables [12]. The parameters obtained from the Tait equation were in an excellent agreement with those using densities from the NIST tables, and averaged values are listed in Table III.

Experimental viscosities of the nine refrigerants were chosen at $T^*=0.5$ - 1.2, corresponding to T/T_c =0.39 to 0.94. (T_c /(ϵ_{LJ}/k) \approx 1.28). The values in Table VI were used together with Eq.(13) and Eq.(14) to obtain the characteristic volume V_o for each refrigerant at a given temperature. Experimental viscosity data were then used to obtain the coupling parameter R_η for each refrigerant. It was found that R_η is constant for R11, R12 and R22, a linear function of temperature for R32, but a quadratic function of

temperature for the ethane based refrigerants. The results are given in Table III.

Similarly, it was found that R_{λ} is a linear function for R32 and ethane based refrigerants, but a quadratic function for R11, R12 and R22. Table III lists the results from the various methods. In general, the results from the different methods are comparable.

The coupling parameters were correlated as follows:

$$R_n = D_1 + D_2 T^* + D_3 T^{*2}$$
 (15)

$$R_{\lambda} = E_1 + E_2 T^* + E_3 T^{*2} \tag{16}$$

where the coefficients D_i and E_i are listed in Table III.

Results for refrigerant mixtures using the mixing rule given in Eqs (9-11) are presented in Table IV

7. Application to Halogenated Methanes

Five halogenated methanes were chosen to further test the RHS3 method. Dymond [13] found that for the halogenated methanes, the translational-rotational coupling is in the order: R10 > R11 > R13 > R14. Therefore, it is important to test whether any of the methods described above can predict such a trend.

LJ parameters and average values of R_η and R_λ were obtained as described above and are given in Table V. The diameter σ_{LJ} decreases by 0.20 (Å) for each substitution from R10 to R14 except between R13 and R14, whereas ϵ_{LJ}/k changes by -60 K with each substitution, and T_c /(ϵ_{LJ}/k) = 1.28-1.26. In the case of the viscosity, R_η shows a good trend with each substitution from Cl to F except the last one (from R13 to R14). This may be because of experimental uncertainty. In the case of the thermal conductivity, the method exhibits excellent trends.

8. Summary

Two approaches for obtaining the parameters of the RHS method for transport properties of dense fluids are described. The first approach (RHS1) determines the characteristic volume V_o and coupling parameter R_η for each substance from viscosity data. R_λ is then obtained from thermal conductivity data using the same value of the characteristic volume. The second approach (RHS2) determines V_o from the effective

hard sphere diameter of Lennard-Jones fluids and the Heyes relationship between the Lennard-Jones diameter and the hard sphere diameter. The LJ parameters required in the calculations were determined from saturated densities at T/T_c = 0.75. The coupling parameters were found to be independent of pressure for both methods. The RHS2 method has the advantages of simplicity, and of yielding parameters which exhibit the correct trends.

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REFERENCES

- 1. M. J. Assael, J. H. Dymond, M. Papadaki, and P. M. Patterson, *Int. J. Thermophys.* **13:**269 (1992).
- 2. M. J. Assael, J. H. Dymond, M. Papadaki, and P. M. Patterson, *Int. J. Thermophys.*, **13:**359 (1992).
- 3. M. J. Assael, J. H. Dymond, M. Papadaki, and P. M. Patterson, *Int. J. Thermophys.*, **13:**895 (1992).
- 4. M. J. Assael, J. H. Dymond, and S. K. Polimatidou, *Int. J. Thermophys.*, **15:**189 (1994).
- 5. M. J. Assael, J. H. Dymond, and S. K. Polimatidou, *Int. J. Thermophys.*, **16:**761 (1995).
- 6. T. F. Sun, J. Bleazard, and A. S. Teja, J. Phys. Chem., 98: 1306 (1994).
- 7. D. Chandler, J. Chem. Phys., **62**: 1358 (1975).
- 8. J. H. Dymond and M. A. Awan, *Int. J. Thermophys.*, **10:**941 (1989).
- 9. J. H. Dymond, Q. Rev. Chem. Soc., 3: 317 (1985).
- 10. D. M. Heyes, J. Chem. Soc., Faraday Trans. 2, **84(6):** 705 (1988).
- 11. T. F. Sun, and A.S. Teja, *J. Phys. Chem.*, **100:** 17365 (1996).
- 12. M. Huber, J. Gallagher, M. McLinden, and G. Morrison, Standard Reference Database
- 23, NIST thermodynamic properties of refrigerants and refrigerant mixture database (REFPROP), version 5.0 Gaithersburg, Maryland: Standard Reference Data Program, National Institute of Stanfards and Technology.
- 13. M. J. Assael, S. K. Polimatidou, E. Vogel, and W. A. Wakeham, *Int. J. Thermophys.* **15:**575 (1994).
- 14. M. J. Assael, J. H. Dymond, and S. K. Polimatidou, *Int. J. Thermophys.*, **15:**591 (1994).
- 15. M. J. Assael and S. K. Polimatidou, *Int. J. Thermophys.*, **15:**779 (1994).
- 16. C. M. B. P. Oliveira and W. A. Wakeham, Int. J. Thermophys. 14:33 (1993).
- 17. C. M. B. P. Oliveira and W. A. Wakeham, *Int. J. Thermophys.* **14:**1131 (1993).
- 18. P. S. van der Gulik, Int. J. Thermophys. 14:851 (1993).

- 19. T. Okubo, T. Hasuo, and A. Nagashima, Int. J. Thermophys. 13:931 (1993).
- 20. A. Kumagai and S. Takahashi, *Int. J. Thermophys.* **12:**105 (1991).
- 21. A. Kumagai and S. Takahashi, *Int. J. Thermophys.* **14:**339 (1993).
- 22. D. Ripple and O. Matar, J. Chem. Eng. Data 38:560 (1993).
- 23. D. E. Diller, A. S. Aragon, and A. Laesecke, Fluid Phase Equil. 88:251 (1993).
- 24. D. E. Diller, S. M. Peterson, *Int. J. Thermophys.* **14:**55 (1993).
- 25. M. J. Assael, L. Karagiannidis, and W. A. Wakeham, *Int. J. Thermophys.* 13:735 (1992).
- 26. J. Yata, T. Minamiyama, and S. Tanaka, Int. J. Thermophys. 5:209 (1984).
- 27. N. Kitazawa and A. Nagashima, Bull. JSME 24:374 (1981).
- 28. M. J. Assael and L. Karagiannidis, Int. J. Thermophys. 14:183 (1993).
- 29. S. H. Kim, D. S. Kim, M. S. Kim, and S. T. Ro, Int. J. Thermophys. 14:937 (1993).
- 30. M. Papadaki and W. A. Wakeham, Int. J. Thermophys. 14:1215 (1993).
- 31. J. Yata, M. Hori, T. Kurahashi, and T. Minamiyama, *Fluid Phase Equil.* **80:**287 (1992).
- 32. M. Papadaki, M. Schmitt, A. Seitz, K. Stefan, B. Taxis, and W. A. Wakeham, *Int. J. Thermophys.* **14:**173 (1993).
- 33. A. Laesecke, R. A. Perkins, and C. A. Nieto de Castro, *Fluid Phase Equil.* **80:**263 (1992).
- 34. U. Gross, Y. W. Song, and E. Hahne, *Int. J. Thermophys.* **13:**957 (1992).
- 35. U. Gross, Y. W. Song, and E. Hahne, *Fluid Phase Equil.* **76:**273 (1992).
- 36. J. Yata, M. Hori, K. Kobayashi, and T. Minamiyama, Presented at 12th STP, Boulder, CO (1994).
- 37. S. T. Ro, J. Y. Kim, and D. S. Kim, Presented at 12th STP, Boulder, CO (1994).
- 38. M. J. Assael and L. Karagiannidis, *Int. J. Thermophys.* **16:**851 (1995).
- 39. M. Arnemann, and H. Kruse, Proc. XVIII th Int. Congr. of Refr., Montreal, Canada, Vol. II. pp 379-383.
- 40. Y. A. Mikhno and V. Z. Geller, Heat Transfer Soviet Research, 16:135 (1984).
- 41. S. T. Ro, J. Y. Kim, and D. S. Kim, Int. J. Thermophys. 16:1193 (1995).

Table I. Comparison of Calculated and Experimental Viscosities

			R	HS3		F	RHS1		F	RHS2		
	Data	a pts	AA	D%	MAD%	6 AA	AD%	MAD9	% AA	AD%	MAD%	6
Liq.	HP	LP	HP	LP		HP	LP		HP	LP		Literature
R11	36	9	0.84	1.15	2.07	0.52	0.73	1.33	0.64	0.64	0.94	13,20
R12	36	8	1.48	2.94	5.08	1.75	2.21	5.74	2.02	4.18	9.87	13,20
R22	43	6	0.85	2.67	6.67	0.74	2.29	5.60	0.85	2.72	6.83	15,20
R32	46	10	3.04	1.51	15.60	1.01	1.19	2.91	1.16	1.42	3.17	14,22,17
R124	124	39	4.17	4.43	41.84	2.92	2.00	12.13	2.96	3.74	14.61	15,22,24
R125	89	44	2.66	2.15	14.45	1.44	1.12	9.21	2.64	1.61	10.72	15,17,22,25
R134a	148	52	2.44	4.07	17.03	3.41	2.36	16.67	3.75	2.15	17.40	14,16,19,23
R141b	61	28	1.71	2.38	7.38	1.92	1.78	9.85	2.30	3.32	11.69	13,21,23
R152a	65	8	1.10	1.84	5.02	0.77	1.83	4.80	0.92	1.99	4.28	13,20,18
Averag	e		2.03	2.57		1.61	1.72		1.92	2.42		

LP= Saturation values

HP= Pressures above saturation

Table II. Comparison of Calculated and Experimental Thermal Conductivities

			R	HS3		R	HS1		RF	HS2		
	Data	a pts	AA	.D%	MAD%	AA	D%	MAD%	AA	D%	MAD%	6
Liq.	HP	LP	HP	LP		HP	LP		HP	LP		Literature
R11	57	11	1.26	2.56	6.11	0.78	1.91	3.48	0.77	1.92	3.45	25-27
R12	40	10	0.78	6.58	28.42	1.16	2.12	3.62	1.51	2.75	5.02	25,26
R22	62	5	2.17	1.06	5.56	0.53	1.54	2.56	0.52	1.89	3.06	26,28,29
R32	55	10	2.78	4.31	8.13	2.82	3.70	6.49	2.85	3.83	6.78	30,36-38
R124	65	8	1.79	2.25	6.74	1.76	1.98	8.57	1.59	1.82	7.42	31,38
R125	24	10	2.77	5.53	16.20	2.33	3.06	8.95	2.33	3.01	8.02	30,38
R134a	113	9	2.31	3.52	9.48	2.33	1.75	7.43	2.26	1.71	7.21	32-34,37
R141b	70	6	1.64	1.68	11.74	1.46	1.03	9.64	1.44	1.02	8.26	31,32,38
R152a	62	5	3.36	1.01	10.98	2.64	1.98	10.96	2.64	1.77	10.39	29,31,34,35
Averag	e		2.10	3.16		1.76	2.12		1.77	2.19		

LP= Saturation values

HP= Pressures above saturation

Table III. Coefficients of eqns 7-8 and eqns 14-16.

Liq.	C_{η}	A_1	$A_2 \times 10^2$	$A_3 \times 10^4$	B_1	$B_2 \times 10^2$	$B_3 \times 10^4$	$\sigma_{\!\scriptscriptstyle LJ}$	ϵ_{LJ}/k
R11	1 157	64.2171	2 0060		1 0126	0.2555	0.0652	4 0779	269.22
R12		62.2981				0.4021			301.21
R22	1.217	42.8546	-3.2715		1.0615	0.2142		4.3388	288.38
R32	1.081	28.7302	-1.8940		1.0654	0.2462		3.8432	270.62
R124	1.085	65.4159	-3.3840		1.2101	0.1284		4.9374	309.92
R125	1.116	55.6437	-3.1591		3.4303	-1.5934	0.3499	4.6831	265.77
R134a	1.248	53.7759	-3.9235		0.7265	0.3496		4.5826	292.12
R141b	0.994	75.9654	-8.9035	1.0770	1.6299	-0.0561		5.0214	374.00
R152a	1.097	58.6224	-11.664	1.4562	0.7585	0.2779		4.4088	300.77
T:-	D	Ъ	D	Г	Е	Г			
Liq.	\mathbf{D}_1	D_2	D ₃	E ₁	. L 2	2 E	3		
D11	1 1777	E		1.7	12770 0	70754	0.72020		
R11	1.1777					.70754			
R12	1.1084	1		2.1	18566 -1	.74739	1.17223		
R22	1.0579	3		1.9	90104 -1	.23951	0.86858		
R32	0.8519	5 0.1273	34	0.9	98171 0	.65083			
R124	4.9254	7 -6.5795	59 2.951	80 1.4	19591 0	.29324			
R125	4.2078	4 -4.7469	5 1.928	09 1.2	25585 0	.64916			
R134a	4.8099	5 -6.3937	73 2.906	35 1.0)9901 0	.71446			
R141b	5.8254	4 -10.858	31 6.395	52 1.6	59667 -0	.12541			
R152a	3.2730	8 -4.1826	52 1.988	05 0.9	93081 0	.66030			

Table IV. Comparison of Viscosities and Thermal Conductivities for Refrigerant Mixtures

Mixtures	P	Data	RH:		RHS		RHS	MAD%	Literature
				Vis	scositiy				
R125a+R134a	LP	15	1.55	2.95	3.84	5.11	3.00	4.56	32
R32+R134a	LP	15	8.87	12.56	7.29	10.20	6.78	10.53	32
R32+R124	LP	14	13.20	18.55	15.62	20.61	14.94	20.11	32
R22+R152a	LP	29	3.29	10.07	2.76	9.17	2.74	9.98	39 ^a
			-	Thermal	Conduct	ivity			
R22+R152a	HP	125	2.20	4.84	2.51	4.89	2.26	5.30	29 a
R12+R22	LP	24	5.69	30.72	2.01	5.52	2.63	6.47	40 a
R32+R134a	HP	120	2.54	10.86	2.72	6.60	2.17	5.78	41 ^a

 $[^]a$ Mixure densities were estimated by x_1 ρ_1+x_2 ρ_2 where x is the mole fractiobn.

Table V. Parameters for Halogenated Methanes

				RHS	RHS2		S3	RH		
Liq.	$T_{c}(K)$	$\sigma_{LJ}(\mathring{A})$	ε_{LJ}/k (K)	R_{η}^{a}	$R_^a$	R_{η}	$R_^a$	R_{η}	$R_^a$	$R_d^{\ b}$
R10	556.35	5.1654	434.85	1.32	1.77					0.54
R11	471.20	4.9768	368.03	1.20	1.67	1.0	1.5	1.2	1.6	0.64
R12	384.95	4.7593	301.16	1.13	1.63	1.6	2.0	1.7	2.1	
R13	302.01	4.5028	237.39	1.08	1.58					0.9
R14	227.50	4.1776	179.92	1.11	1.52					1.0

 $^{^{\}text{a}}$ R_{η} and R_{η} are averaged values.

^b R_D From Dymond (1985)